

[54] **CATIONIC EMULSIONS OF BITUMINOUS BINDERS OF THE TYPE BITUMEN/POLYMER AND CATIONIC EMULSIFYING SYSTEM PREFERABLY UTILIZABLE FOR OBTAINING SAID EMULSIONS**

[75] **Inventors:** Francis Demangeon, Dardilly; Emile Lopez, Vienne, both of France

[73] **Assignee:** ELF France, Paris, France

[*] **Notice:** The portion of the term of this patent subsequent to Sep. 20, 2005 has been disclaimed.

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[30] **Foreign Application Priority Data**

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[58] **Field of Search** 524/61, 60, 68, 59; 252/311.5

[56] **References Cited**

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Primary Examiner—Allan M. Lieberman

Attorney, Agent, or Firm—Burgess, Ryan & Wayne

[57] **ABSTRACT**

This invention provides cationic emulsions of bituminous binders of the bitumen/polymer type, formed by a dispersion of an organic phase of the binder in an aqueous phase containing a cationic nitrogenous emulsifier and a mineral or organic acid, the acid being in appropriate quantity so that the pH of the aqueous phase is comprised between 3 and 9. The aqueous phase also contains an agent consisting of at least one water-soluble sequestering salt of phosphonic acid, particularly a salt of alkaline metal or of amine of hydroxyalkyl-1,1-diphosphonic acid, and optionally a water-soluble thickening agent. To obtain the emulsions, which are usable for road applications, the emulsifying agent, the sequestering agent and optionally the thickening agent may be used in the form of a premixture containing the agents and called cationic emulsifying system.

21 Claims, No Drawings

**CATIONIC EMULSIONS OF BITUMINOUS
BINDERS OF THE TYPE BITUMEN/POLYMER
AND CATIONIC EMULSIFYING SYSTEM
PREFERABLY UTILIZABLE FOR OBTAINING
SAID EMULSIONS**

This is a division of application Ser. No. 932, 502, filed as PCT FR86/00040 on Feb. 12, 1986, published as WO86/04915 on Aug. 28, 1986, now U.S. Pat. No. 4,772,648.

FIELD OF THE INVENTION

The invention relates to cationic emulsions of bituminous binders of the bitumen/polymer type. In one aspect, the invention relates to a cationic emulsifying system preferably utilizable for obtaining these emulsions.

BACKGROUND OF THE INVENTION

Converting bituminous binders in cationic or anionic emulsions is a well-known technique. The emulsion consists of a dispersion of fine bitumen globules in a continuous aqueous phase. The emulsion is called anionic or cationic according to the total electrical charge surrounding the globule which can be negative or positive. In the presence of a continuous electric current, the micellae of bitumen of an emulsion settle on the anode if the emulsion is anionic or on the cathode if the emulsion is cationic.

The conversion of bituminous binders to emulsion makes use of emulsifying agents that favor the dispersion of the globules of binder in the aqueous phase by forming a charged protective shell that prevents their agglomeration.

In the case of anionic emulsions, the emulsifying agent is of the anionic type and preferably consists of a fatty acid alkaline salt. This emulsifying agent is generally mixed with the bituminous binder in a quantity of from 0.05 to 2% by weight. The binder containing the emulsifying agent is then dispersed in an alkaline aqueous phase by passing into an emulsifier.

Cationic emulsions are obtained by using cationic emulsifying agents, which generally consist of nitrogenized organic compounds such as fatty amines, fatty amidoamines, fatty imidoazolines, fatty mono and diquaternary ammonium compounds and reaction products of one or more of said products with ethylene or propylene oxide. The fatty radical of the emulsifying compounds can possess a very diversified chemical structure and the raw materials for the preparation of the emulsifying compounds can originate from different sources such as animal fats, refined petroleum products, vegetable oils and liquid resins. The cationic emulsifying agent is generally dispersed in water to which hydrochloric acid is added. The hydrochloride formed is soluble. The homogeneous solution thus obtained, having a pH generally comprised between 1 and 5, is then put into contact with the bituminous binder in an emulsifier in order to form the emulsion.

Regardless of whether it is anionic or cationic, the emulsion is considered to be a means that allows lowering the viscosity of bituminous binders. The emulsion restores, after rupture, the bituminous binder to which has been added the emulsifying agent used for producing the emulsion. The presence of this emulsifying agent in the bituminous binder is regarded as exerting a favorable effect on the adhesiveness of the binder in the presence of mineral surface, provided it is within a narrow range of pH comprised between 1 and 6.

The largest sale of bituminous emulsions is mainly in the area of building and repair of roads, and essentially in the form of cationic emulsions. Experience in this field shows that cationic emulsions are easier to apply and prevent the numerous inconveniences of anionic emulsions such as the possibility of restoration to emulsion in the presence of water, of unforeseeable rupture in storage and of weak adhesiveness of the binder to the surface of the aggregate.

Cationic bituminous emulsions are mainly used according to two type of technique, namely, coating and encasing.

Coating consists in gluing layers of fine gravel on the surface of the road by means of the bituminous emulsion in order to produce roughness and make the surface impervious. This technique contributes to the creation of a new runway layer.

In encasement, the bituminous emulsion allows the use of layers having a thickness of from 0.4 cm to more than 20 cm, which integrate in all levels of a road structure.

At present, due to the general reduction of budgets for repair and building of roads, the use of the bitumen emulsion is particularly limited to the repair of the surface of roads. However, due to the extreme mechanical demands to which are subjected the mosaics of the surface coatings, the road bitumens of emulsifiable kinds do not have the characteristics necessary that will allow them to bear the tangential stresses due to heavy loads and heavy traffic.

Under these circumstances, the bituminous binders exclusively constituted of bitumen are being increasingly replaced by bituminous binders of the bitumen/polymer type. The bitumen/polymer binders are products obtained from bitumens to which polymers have been added and have eventually been modified by the latter. Optionally, a reagent such as sulfur or a modifying agent and/or an adjuvant can also be present.

Examples of bituminous binders of the bitumen/polymer type comprise preferably:

bitumens modified by olefinic polymers or by mixtures of fatty acids and higher alcohols (French Patent No. 7,627,936 publication No. 2,364,960] of Sept. 17, 1976);

bitumens modified as above and on which have been grafted elastomers (French Patent No. 7,627,937 [publication No. 2,364,961] of Sept. 17, 1976);

bitumens modified by norbornene (French Patent No. 7,630,316 [publication No. 2,367,102] of Oct. 8, 1976);

bitumens modified by dihalopolybutadienes serving as receptive structures to block copolymers and to covulcanizable elastomers (French Patent No. 7,806,160 [publication No. 2,418,812] of Mar. 3, 1978);

bitumens modified by block copolymers styrene/butadiene or styrene/isoprene (French Patent No. 7,639,233 [publication No. 2,376,188] of Dec. 28, 1976);

bitumens modified by block copolymers styrene/carboxylated diene (addition No. 7,831,689 [publication No. 2,440,967] of Nov. 9, 1978 to French Patent No. 7,639,233);

bitumens modified by polyethylene waxes (French Patent No.

[publication No. 2,405,288] of Oct. 5, 1977);

bi-tumens modified by elastomeric acrylic polymers (French Patent No. 7,812,135 [publication No. 2,424,301] of Apr. 25, 1978;

bitumens modified by polymers of the type polyvinyl alcohol, polyamide, polyester, polyurethane;

bitumens modified by olefinic copolymers, preferably among them, ethylene/propylene copolymers or ethylene/propylene/diene copolymers, or by copolymers of olefins and of vinyl monomers, preferably ethylene/vinyl acetate copolymers, ethylene/acrylic ester copolymers, ethylene/vinyl chloride copolymers;

bitumens modified by special polymers of the type of halogenated polymers and preferably fluorinated polymers such as poly-tetrafluoroethylenes or fluorinated and chlorinated polymers, or of the type of silicon polymers such as polysiloxanes;

bituminous compositions obtained by bringing into contact one bitumen with a mother solution containing one polymer, preferably one block copolymer styrene/-conjugated diene and sulfur or a polysulfide or another reactive agent, preferably vinyl monomer, as well as a petroleum fraction and/or a coal oil (French Patents No. 7,818,534 [publication No. 2,429,241] of June 21, 1978 and No. 8,210,095 [publication No. 2,528,439] of June 10, 1982; French Patents No. 7,910,987 [publication No. 2,455,623] of May 2, 1979 and No. 8,216,433 [publication No. 2,533,935] of Sept. 30, 1982);

bitumens modified by a mixture of at least two of the polymers cited above, and mixtures of said modified bitumens.

However, the emulsification of said different bitumens modified by the incorporation of polymers cannot be effected as such, since the improvement on the properties of the bitumens systematically increases the viscosity of the medium, which hinders the preparation of binders modified by polymers making use of the conventional industrial equipments. Under these circumstances, it is necessary to use one or several fluidizing solvents, the main part they play being to reduce the viscosity of the medium and bring it to a range equivalent to that of operation of bitumens of penetration 80/100 and 180/220 between 100° C. and 150° C., that is, 1 to 10 poises. The solvent or solvents selected have among other properties that of being partially volatile while preserving the heavy fractions of plastifying and compatibilizing properties between the polymer and the bitumen.

The applicant has observed that the formulation of emulsions of quick rupture from binders of the bitumen/polymer type fluidized or fluxed by oils of petroleum origin or resulting from the distillation of coal in the presence of conventional cationic emulsifying agents such as those mentioned above, led to emulsions in which the rupture is poor or uncertain. It has been observed, on the other hand, that this rupture was represented by a partial coagulation of the bitumen globules without ever resulting in the first hours, even after several days, in the total coalescence of the micellae of the emulsion, and this, whatever the type of emulsifying agent used and the concentration of said emulsifying agent. This deficiency confers on the residual bituminous binder mechanical characteristics which are very inferior to those of the original anhydrous bitumen/polymer binder. Therefore, it limits the development of the emulsions of bitumen./polymer binder in the field of producing thin and superficial coatings on roads having very heavy traffic. The applicant has demonstrated in addition that the phenomenon exclusively depended on a double interaction between the solvent and the emulsifying agent for a conventional emulsifying agent such as defined above and between the solvent and the salts

contained in the water, namely, alkaline earth salts and metallic salts. The nature of the bitumen and its origin, the type of polymer associated with the bitumen, and the presence of cross-linking or stabilizing reagent, on the contrary, exert no influence on the phenomenon.

SUMMARY OF THE INVENTION

It has been found that the deleterious influence of the interactions between solvent and emulsifying agent and between solvent and salts contained in the water could be suppressed by using in the cationic emulsion of bitumen/polymer binder a specific emulsifying system that results from the association of at least one cationic nitrogenized emulsifying agent with an adjuvant consisting of at least one water-soluble sequestering salt of phosphonic acid.

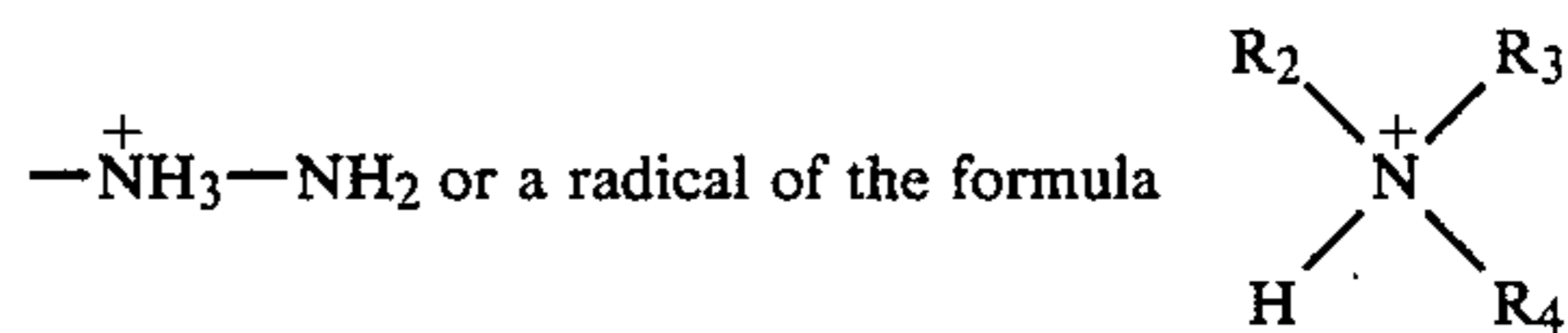
Therefore, an object of the invention is to provide cationic emulsions of bituminous binders of the bitumen/polymer type that are formed in a dispersion of an organic phase consisting of the bitumen/polymer binder in an aqueous phase including a cationic nitrogenized emulsifying agent and an acid, the latter being present in an amount such that the pH of the aqueous phase has a value comprised between 3 and 9, the emulsions being characterized by the fact that their aqueous phase also includes an adjuvant consisting of at least one water-soluble sequestration salt of phosphonic acid in which the sequestering power in relation to the metallic ions is equivalent, in a pH range of from 3 to 9, to a fixation of at least 5 mg calcium ions per gram of salt.

DETAILED DESCRIPTION OF THE INVENTION

The water-soluble sequestering phosphonic salt or salts are preferably selected among the diphosphonic salts of the formula (I)



wherein R₁ is a hydrocarbyl radical of C₁ to C₁₀, preferably an alkyl, cycloalkyl or aryl radical of C₁ to C₆, and the M symbols, which can be identical or different, each designate a hydrogen atom or an M₁ group in a manner such that at least one M is an M₁ group the M₁ group representing one alkaline metal atom, the radical



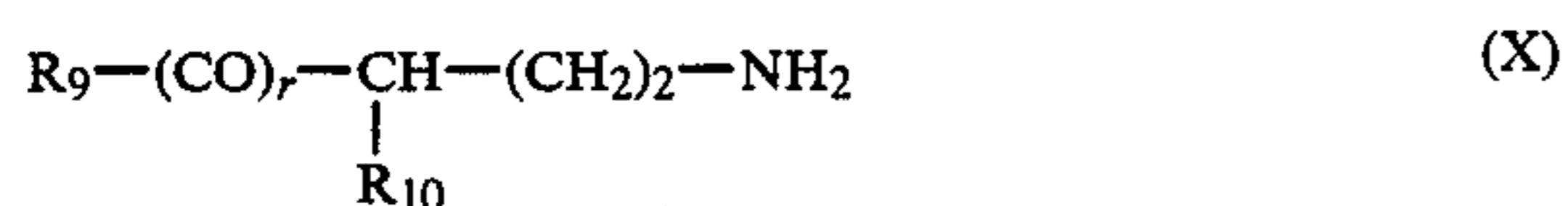
wherein the symbols R₂ to R₄, which can be identical or different, each represents one hydrogen atom or an organic radical and in particular a hydrocarbyl monovalent radical of C₁ to C₈, preferably C₁ to C₆, and which may include one or several functional groups and particularly an OH group, some of the R₂ to R₄ radicals can be combined to form a heterocycle with the nitrogen atom to which they are bonded. The salts themselves are not emulsifying agents.

The alkaline metal that can represent the M₁ group is selected among the metals of group IA of the Periodic Classification of Elements such as shown in the HANDBOOK OF CHEMISTRY AND PHYSICS, 46th Edi-



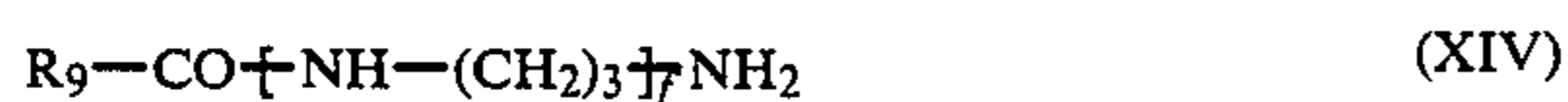
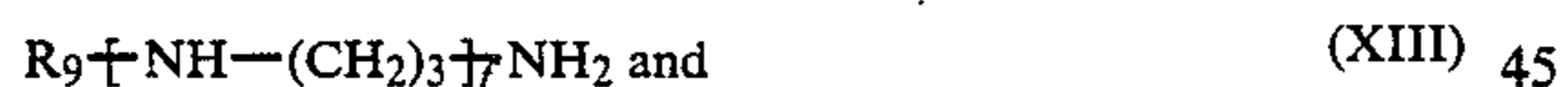
s representing in these formulae an integer from 1 to 6, preferably from 1 to 3, the R₇'s, which can be identical or different, designate C₆ to C₂₂ hydrocarbonyl radicals which can be substituted with one or several functional groups and preferably alkyl or alkenyl radicals from C₈ to C₂₂ or a benzyl radical, the R₈'s, identical or different, representing alkyl radicals from C₁ to C₆, which can be hydroxylated, preferably methyl, ethyl, propyl, hydroxyethyl, hydroxypropyl, Y⁻ designating an anion of a mineral acid, in particular a chloride anion, or of an organic acid, preferably an acetate or format anion, h being a number equal to (4 - j) and j being a number assuming the values 1, 2 or 3, while R, r, m, n and q have the meanings given above.

The preferred cationic nitrogenized emulsifying agents that can be used as emulsifying agents A have the following formulae



in these formulae an alkyl or alkenyl radical from C₁₂ to C₂₂, preferably stearyl and/or oleyl, R₁₀ designating an alkyl or hydroxyalkyl radical from C₁ to C₃, Y⁻ being a chloride or acetate anion, 1 being a number equal to (3 - g) and g being a number assuming the values 1, 2 or 3 while r, j and h are numbers having the meanings given above.

Other preferred cationic nitrogenized emulsifying agents that can be preferably used as emulsifying compounds B correspond to the following formulae (XIII) and (XIV):



wherein f designates a number assuming the values 2, 3, 4 or 5 and R₉ has the above meaning.

The total quantity of cationic nitrogenized emulsifying agent in the emulsion can change widely. This quantity can advantageously represent from 0.03% to 3%, preferably from 0.06% to 2%, of the total weight of the emulsion for obtaining emulsions of quick rupture and satisfactory stability.

The acids used for adjusting the pH of the emulsion to a value of from 3 to 9 can be selected among the mineral acids of different degrees of acidity or also among the mono- or polycarboxylic acids, saturated or unsaturated, and eventually hydroxylated which are defined in French Pat. No. 8,122,275 (publication No. 2,517,317) of Nov. 27, 1981. Preferred acids are hydrochloric, nitric, phosphoric, acetic, oxalic, citric and formic acids.

The bitumen-polymer binder reduced to cationic emulsion can be any one of the products obtained from bitumens to which have been added one or several

polymers and eventually modified by this or these polymer(s) in the presence or absence of a reagent such as sulfur or an agent that releases sulfur such as a polysulfide, of a coupling or vulcanization agent, it being likewise possible to add to said bitumens, if necessary, a fluxing agent such as a petroleum fraction, a coal oil, or a mixture of said products.

Preferred bitumen/polymer binders can be selected among the different bitumen/polymer binders cited above by way of example.

The emulsion according to the invention can also include a thickening agent consisting of one or several water-soluble organic compounds, which, within a pH range of from 3 to 9, are capable of increasing the viscosity of the emulsion without modifying the viscosity of the aqueous phase alone containing the cationic nitrogenized emulsifying agent.

Among the compounds that can be used as thickening agents, there can preferably be mentioned the water-soluble natural gums of the type guar gum, arabic gum, ghatti gum, karaya gum, tragacanth gum, locust bean gum, or also the water-soluble polyurethanes of low molecular weight, and in particular polyurethanes of molecular weight below 20,000 which result from the reaction of a polyisocyanate such as toluene diisocyanate, diphenyl methane 4,4'-diisocyanate, naphthylene 1,5-diisocyanate, hexamethylene 1,6-diisocyanate, 3,3'-tolidene - 4,4'-diisocyanate and m-phenylene disulphonyl diisocyanate with polyols of the polyester polyol or polyether polyol type. As polyester polyols can be mentioned the reaction products of a polycarboxylic acid such as adipic, phthalic or maleic acid with a polyol or a mixture of polyols such as ethylene glycol, propylene glycol, diethylene glycol, butanediol, glycerol, trimethylol propane, hexamethylene triol and pentaerythritol, while appropriate polyether polyols can be selected among the reaction products of polyols of the ethylene glycol, propylene glycol, glycerol, trimethylolpropane, pentaerythritol, methyl glycoside, sorbitol and sucrose type with epoxides of the ethylene oxide, propylene oxide and epoxystyrene type.

The weight of thickening agent in the emulsion can advantageously represent from 5% to 30% of the total weight of the combination formed by the cationic nitrogenized emulsifying agent, the phosphonic adjuvant and the thickening agent.

The cationic emulsions according to the invention are prepared by making, in a zone of emulsion formation, particularly in an apparatus of the emulsifier type, a homogeneous dispersion of the bitumen/polymer binder in an aqueous phase containing, in quantities adequate for obtaining the concentrations defined above, the emulsifying agent salified by an acid in an amount sufficient for obtaining the desired pH value, the water-soluble phosphonic sequestering agent and optionally the thickening agent. To do this, there are separately and simultaneously brought to the zone of formation of the emulsion, on one side, the bitumen/polymer binder in the form of a molten mass having a temperature between 80° C. and 180° C., preferably between 120° C. and 160° C., and on the other hand, the aqueous phase containing the above mentioned ingredients and preferably between 20° C. and 60° C. and the combination is maintained in said zone for a time sufficient for forming an emulsion.

The aqueous phase, which is mixed with the bitumen/polymer binder in the zone of formation of the

emulsion, is prepared by introducing the emulsifying agent, the phosphonic sequestering compound, the acid and the eventual thickening agent in the amount of water necessary for producing the emulsion, which has been previously brought to a temperature comprised between 30 and 75° C. The quantities of emulsifying agent, of phosphonic sequestering compound and of eventual thickening agent which are added to the water are selected in a manner such that the concentrations of said ingredients in the resulting emulsion be within the above defined ranges. The acid is used in a quantity sufficient for transforming, if necessary, the cationic emulsifying agent into a soluble salt and for bring the pH of the emulsion to the selected value. The ingredients, namely, cationic nitrogenized emulsifying agent, phosphonic sequestering compound, acid and thickening agent, can be added to the water in any order. The phosphonic sequestering compound in particular can be introduced in the water before or after salification of the emulsifying agent by the acid. The phosphonic sequestering compound can likewise be formed in situ in the water from the corresponding precursor phosphonic acid and from a base apt to furnish alifying groups M₁ such as defined above.

In an advantageous form of preparation, at least the emulsifying agent and the phosphonic sequestering compound, or the phosphonic acid precursor thereof, are simultaneously incorporated into the water in the form of a mixture called cationic emulsifying system which has been previously made, the acid is then added and the thickening agent if it is used.

The cationic emulsifying system includes, therefore, a cationic nitrogenized emulsifying agent consisting of one or more cationic nitrogenized emulsifying agents and an adjuvant consisting of at least one water-soluble phosphonic sequestering compound such as defined above or the phosphonic acid precursor thereof, said adjuvant being in particular formed of at least one phosphonic compound selected among the compounds of formula (I) or the phosphonic acids precursors thereof of formula (II).

In such an emulsifying system the weight of the cationic nitrogenized emulsifying agent represents in particular from 10 to 99%, preferably from 25 to 90%, of the total weight of emulsifying agent and adjuvant.

The cationic nitrogenized emulsifying agent present in the cationic emulsifying system can consist preferably of one or more of the cationic nitrogenized emulsifying agents selected among the fatty monoamines of formula (III) and the polyamines and amidopolyamines of formula (IV).

In a preferred manner of preparation, the cationic nitrogenized emulsifying agent of the cationic emulsifying system is constituted by the association of one or more cationic nitrogenized emulsifying agents A with one or more cationic nitrogenized emulsifying agents B such as defined above, the weight ratio of the total amount of the compound or compounds A to the total amount of compounds A and B being in particular from 5 to 95%.

In particular the compounds A can be selected among the compounds defined above by formulae (III), (V), (VI) and (VII) and preferably among the compounds A defined by formulae (IX), (X), (XI) and (XII), while the B compounds can be selected among the compounds defined above by the formula (VIII) and preferably among the compounds B defined by the formulae (XIII) and (XIV).

The cationic emulsifying system can also contain a thickening agent consisting of one or more water-soluble organic compounds selected among those defined above, and in particular among the water-soluble natural gums or among the water-soluble polyurethanes of low molecular weight and preferably those resulting from the reaction between a polyisocyanate and a polyol of the polyester polyol or polyether polyol type.

The weight of thickening agent in the cationic emulsifying system advantageously represents from 5% to 30% of the total weight of the mixture comprising the emulsifying agent, the phosphonic adjuvant and the thickening agent.

In the preparation of the emulsion, the aqueous phase containing the cationic nitrogenized emulsifying agent, the phosphonic adjuvant, the acid and the thickening agent, on one hand, and the bitumen/polymer binder, on the other, are brought to the zone of formation of emulsion in proportions such that the cationic emulsion resulting advantageously contains by weight from 30 to 85%, preferably from 50 to 80%, of bitumen/polymer binder.

The cationic emulsions of bitumen/polymer binder according to the invention are characterized by a quick and irreversible rupture upon contact with the aggregates at the end of which the bitumen/polymer binder immediately and entirely recovers its initial properties.

The emulsions according to the invention are particularly suited to applications of road building and repair, to caulking in civil engineering or in construction and also to industrial uses.

The invention is illustrated by the examples that follow.

In these examples, which describe cationic emulsions of bituminous binders prepared according to the prior art (control examples) and according to the invention, the quantities appearing below are given to characterize the samples of emulsion prepared:

medium diameter of the globular distribution of the bituminous binder, that is, the value of the diameter corresponding to the value of 50% on the curve of cumulative gravimetric particle-size distribution of the globules of the bituminous binder obtained by laser granulometric analysis;

index of rupture in fine siliceous product (abbreviated IR) determined according to the French standard NF T 66 017 and representing the minimum quantity, expressed in grams of siliceous sand of granulometry from 40 to 150 μ m, entailing the rupture of 100 g of emulsion with formation of a bituminous clot sand/binder (a value of the rupture index of about 80 corresponding to a quick rupture of the emulsion, which is what is sought);

time of rupture of the emulsion on plates of bituminous coatings with the aid of microdorit fine gravel having a diameter of from 10 to 14 mm (on a surface formed by a bituminous coating the emulsion is placed, thereafter said emulsion is coated by means of fine gravel, and there is counted from the time the emulsion is covered by the fine gravel, the time that elapses until the rupture of the emulsion, which appears by an attachment of the fine gravel to the support).

Besides, the following characteristics have likewise been determined in the course of the evaluation test of the rupture index:

quantity of liquid salted out after obtention of the clot of fine sands coated by the bituminous binder ruptured and content of water of this liquid (a small amount of

liquid containing an elevated amount of water resulting in a quick and open rupture of the emulsion); and

cohesion/elasticity of the clot of fine sands coated by the bituminous binder since the formation of said clot, determined by traction test according to the French standard NF T 46 002.

The composition of the liquid, that is, of the water/emulsion phase salted out during the rupture of the emulsion, objectively shows the quantity of non-ruptured emulsion, this information being related to the phenomenon of delay or arrest in the rupture of the emulsion. A small amount of liquid containing an elevated amount of water corresponds to a quick and open rupture of the emulsion whereas a larger amount of liquid containing a small amount of water shows a slow rupture of the emulsion, which is what has to be prevented.

The immediate cohesiveness or elasticity of the clot of fine siliceous sands coated with bituminous binder is a characteristic that accounts for the coalescence of the mass of globules of the ruptured bituminous binder, and that gives an indication of the capacity of the bituminous binder to recover its initial viscoplastic or elastomeric properties.

The emulsions described in the examples have been prepared using, as the case may be, either city water having a total hardness corresponding to a hydrotimetric standard equal to 22° or an industrial water having a total hardness corresponding to a hydrotimetric standard of 24° and containing 15 mg per liter of ferrous ions.

In the examples given below the quantities and percents are given by weight unless otherwise indicated.

EXAMPLE 1

In this example, which is a control example, there was prepared a cationic emulsion at 70% of a bituminous binder consisting of an ordinary bitumen of 180/220 penetration, as follows.

There are formed in the first place an aqueous phase consisting of an aqueous solution of hydrochloride of the emulsifying agent, producing a 0.16% dispersion of propylene diamine-1,3 of tallow (cationic nitrogenized emulsifying agent) in industrial water, thereafter adding to the dispersion 0.14% hydrochloric acid at 20° Be (d=1.16).

The emulsion was then produced in an emulsifier in which there were simultaneously and separately introduced 100 parts of the aqueous phase prepared as indicated above and brought to 45° C., and 700 parts of the bitumen brought to 145° C.

There were obtained 1000 parts of a control cationic emulsion the characteristics of which are given below:

pH: 3.5

medium diameter of the bitumen globules (μm): 3-5

index of rupture: 70

amount of salted out liquid after 1 hour (% of the emulsion): 6

amount of water in the salted out liquid (%): 95

cohesiveness of the clot: good

rupture time of the emulsion on coated plates (mn): about 30

The amount of salted out liquid is small, and it is essentially water. The time of rupture of the emulsion on coated plate is about 30 minutes for a rupture index of 70. The cationic emulsion thus produced has a behavior deemed satisfactory with regard to the rupture phenomenon.

EXAMPLE 2

In this example, which represents a second control example, there were prepared three cationic emulsions (tests I, II and III) at 70% of a bituminous binder of the bitumen/polymer type constituted by a reaction product at elevated temperature of a road bitumen of 80/100 penetration with a mother solution consisting of a solution of sulfur and a block copolymer of styrene and butadiene, containing 25% styrene and 75% butadiene in a petroleum fraction obtained in refinery after catalytic cracking of the heavy distillates called "Light Cycle Oil", said fraction having a distillation range on the order of from 180° C. to 360° C.

For preparing the bitumen/polymer binder, the work was carried out as follows. Twenty parts of block copolymer were dissolved in 80 parts of the petroleum fraction at a temperature comprised between 80° C. and 100° C. After the polymer was completely dissolved, 0.6 part of sulfur was added to the solution. Fifteen parts of the solution thus were mixed with 85 parts of the road bitumen and the mixture was brought to a temperature comprised between 170° C. and 180° C. for about 1.5 hours. There was thus obtained a bitumen/polymer binder, the main characteristics of which are indicated herebelow.

viscosity at 160° C.: 89 mPa.s

pseudo-viscosity at 50° C. with an orifice of 10 mm

(NF T 66 005): 115 seconds

traction test at -10° C. with a speed of 500 mm/mn

Stress at threshold: $5.7 \times 10^5 \text{ Pa}$

Stress at breakage: $1.07 \times 10^5 \text{ Pa}$

Elongation at threshold: 20%

Elongation at breakage: >900%

The bitumen polymer binder was emulsified by the following procedure.

There are formed in the first place an aqueous phase consisting of an aqueous solution of hydrochloride of the emulsifying agent, producing a dispersion of propylene diamine-1,3 of tallow (cationic nitrogenized emulsifying agent) in industrial water, thereafter adding to the dispersion 0.14% hydrochloric acid at 20° Be.

Percentages in the aqueous phase of the amounts of emulsifier and hydrochloric acid employed, represented respectively,

0.15% and 0.14% for case I; 0.25% and 0.21% for case II and 0.08% and 0.06% for case III.

The emulsion was then produced in an emulsifier in which there were simultaneously and separately introduced 300 parts of the aqueous phase prepared as indicated above and brought to 45° C., and 700 parts of the bitumen brought to 145° C.

There were obtained 1000 parts of control cationic emulsions of which the characteristics are given below:

Test	I	II	III
pH	3.5	4	4.2
medium diameter of the globules of binder (μm)	3-5	3-5	3-6
rupture index	50	80	40
quantity of salted out liquid after 1 hour (% of the emulsion)	35	21	45
quantity of water in the salted out liquid (%)	32	60	32
cohesiveness of the clot	zero	zero	zero
time of rupture of the emulsion	>180	>180	>180

-continued

Test	I	II	III
on plates of coated products (mn)			

in the three tests all the bitumen/polymer binder is emulsified and the cationic emulsions produced have a good stability in storage.

On the other hand, although in the three emulsions the values of the index of rupture were from quick to very quick ruptures, the quantity of salted out liquid in the test for determining said index is considerable and this liquid contains a high rate of non-ruptured emulsions. Besides, the clot of the mixture of sand and bitumen/polymer binder formed after rupture of the emulsion shows no cohesiveness and in addition the time of rupture of the emulsion on plates of coated material is considerable in each case.

EXAMPLES 3 to 5

In these examples cationic emulsions according to the invention were prepared with 70% of a bituminous binder of the bitumen/polymer type identical with the one described in Example 2.

The bitumen/polymer binder was emulsified using the mode of operation that follows.

There was first formed an aqueous phase consisting of an aqueous solution of the hydrochloride of the emulsifying agent by producing a dispersion of the emulsifying agent in industrial water (Example 3) or in city water (Examples 4 and 5), and then adding to the dispersion, counted in percents of the aqueous phase, 0.13% of hydrochloric acid of 20° Be, then 0.2% of an aqueous solution with 40% tetrapotassium salt of hydroxypropane-1,1 diphosphonic acid.

The emulsifying agent used consisted of a mixture of tallow propylene - 1,3 diamine (emulsifying agent of type A) and of a tallow polypropylene polyamine sold under the name of LILAMULS EM30 (emulsifying agent of type B) in quantities respectively representing, in percents of the aqueous phase, 0.015% and 0.135% in Examples 3 and 4, and 0.16% and 0.02% in Example 5. The aqueous phase obtained in each case was limpid.

The emulsion was then produced in an emulsifier in which there were simultaneously and separately introduced 300 parts of the aqueous phase prepared as indicated above and brought to 45° C. and 700 parts of the bitumen/polymer binder brought to 145° C.

In each example there were obtained 1000 parts of cationic emulsions, the characteristics of which are given herebelow.

Example	3	4	5
pH	5	5.2	5.3
medium diameter of the binder globules (μm)	4-7	3-6	3-7
rupture index	60	60	65
amount of salted out liquid after 1 hour (% of the emulsion)	7	7	9
amount of water in the salted out liquid (%)	95	95	97
cohesiveness of the clot	strong elasticity	strong elasticity	strong elasticity
time of rupture of the emulsion on plates of coated products (mn)	30-60	30-60	30-60

For the three examples all the bitumen/polymer binder was emulsified and the stability in storage of the emulsions produced is good.

The rupture indices of the emulsions are low, and there was found a behavior of the emulsions similar to that of an ordinary bitumen emulsion, namely, small amount of salted out liquid phase and small amount of non-ruptured emulsion.

Besides, the clot of sand and bitumen/polymer obtained in the course of the test for determining the rupture index showed a strong cohesiveness-elasticity and can be subjected to a characterization by traction test.

Herebelow are given the traction characteristics of the clot resulting from the rupture of the emulsion of Example 3 in the course of the test for determining the rupture index, the traction test having been carried out according to the standard NF T 46 002, working at 0° C. with a traction speed of 500 mm/mn.

Stress at threshold: 2.55×10^5 Pa

Stress at breakage: 0.68×10^5 Pa

Elongation at threshold: 15%

Elongation at breakage: 570%

EXAMPLE 6

There was prepared a cationic emulsion of a bituminous binder of the bitumen/polymer type obtained as described in Example 2, but replacing the petroleum fraction "Light Cycle Oil" by the same amount of an anthracenic coal oil having a distillation range on the order of from 200° to 400° C.

The emulsion was prepared as follows:

There was first formed an aqueous dispersion of emulsifying agent by dispersing in industrial water a mixture of cationic nitrogenized emulsifying agents consisting of stearic propylene amidoamine (emulsifying agent of type A) and an alkyl amidopolyamine sold under the name of EMULSAMINE L 60 (emulsifying agent of type B), said emulsifying agents being used in amounts respectively representing 0.02% and 0.15% of the aqueous dispersion. There were then introduced in to the dispersion, expressed in percents of the latter, 0.2% of an aqueous solution with 40% of tetrapotassium salt of hydroxypropane-1,1 diphosphonic acid and then 0.25% of an acetic acid solution at 96% (density = 1.06). There was obtained a limpid aqueous phase.

In the emulsifier there was separately and simultaneously introduced 300 parts of the aqueous phase prepared as indicated above and brought to 45° C. and 700 parts of the bitumen/polymer binder brought to 145° C.

There were obtained 1000 parts of cationic emulsion having the characteristics given herebelow.

pH: 5.6

medium diameter of the globules of binder (μm): 3-6

rupture index: 60

quantity of liquid salted out after 1 hour (% of the emulsion): 6

quantity of water in the salted out liquid (%): 98

cohesiveness of the clot: great elasticity

time of rupture of the emulsion on plates of coated product (mn): 30-60

The whole binder is emulsified and the stability in storage of the emulsion produced is good.

The rupture index of the emulsion is low and the rupture of the emulsion is accompanied by the salting out of a reduced amount of liquid containing little of the non-ruptured emulsion. Besides, the clot of sand and bitumen/polymer binder obtained in the course of the

test for determining the index of rupture has a strong cohesiveness-elasticity.

EXAMPLE 7

There was prepared a cationic emulsion of a bituminous binder of the bitumen/polymer type obtained by mixing a road bitumen of 80/100 penetration with a ethylene/vinyl acetate copolymer containing 45% vinyl acetate.

The binder was prepared by dispersing, by stirring and for about 2 hours, 50 parts of the copolymer in 950 parts of the bitumen brought to 180° C., then fluidizing the homogeneous solution obtained by adding 10% of an anthracenic coal oil having a distillation range on the order of from 200 to 300° C.

The main characteristics of the binder obtained are indicated herebelow:

viscosity at 160° C.: 97 mPa.s
 pseudo-viscosity at 50° C. with an orifice of 10 mm (NF T 66 005): 128 seconds
 traction test at -10° C. with a speed of 500 mm/mn
 Stress at threshold: 12.3×10^5 pa
 Stress at breakage: 6.3×10^5 Pa
 Elongation at threshold: 15%
 Elongation at breakage: 320%

The binder was emulsified by following the mode of operation described in Example 6.

There were obtained 1000 parts of cationic emulsion of bitumen/polymer binder having the characteristics given herebelow.

pH: 5.5
 medium diameter of the globules of binder (μ m): 3-6
 rupture index: 58
 quantity of salted cut liquid after 1 hour (% of the emulsion): 6
 quantity of water in the salted out liquid (%): 99
 cohesiveness of the clot: strong cohesiveness
 time of rupture of the emulsion on plates of coated product (mn): 30-60

All of the binder is emulsified during the preparation of the emulsion and the stability in storage of the emulsion produced is good.

The rupture index of the emulsion is low and the rupture of the emulsion is accompanied by the salting out of a reduced amount of liquid containing very little of the non-ruptured emulsion. Additionally, the clot of sand and bitumen/polymer binder obtained in the course of the test for determining the rupture index has a strong cohesiveness.

EXAMPLE 8

There were prepared two cationic emulsions of a bituminous binder of the bitumen/polymer type obtained by mixing a road bitumen of penetration 80/100 with a stellate triblock copolymer styrene/butadiene/styrene sold under the name of SOLPRENE 411 by PHILLIPS PETROLEUM.

The binder was prepared by dispersing, by stirring for 2 hours, three parts of the copolymer in 100 parts of the bitumen brought to 180° C., then fluidizing the homogeneous solution obtained by adding 12% of a petroleum fraction of the "light Cycle Oil" type having a range of distillation of from 180° C. to 360° C.

The main characteristics of the binder obtained are given herebelow:

viscosity at 160° C.: 102 mPa.s
 pseudo-viscosity at 50° C. with an orifice of 10 mm (NF T 66005): 132 seconds

traction test at -10° C. with a speed of 500 mm/mn

Stress at threshold: 7.1×10^5 pa

Stress at breakage: 0.05×10^5 Pa

Elongation at threshold: 20%

Elongation at breakage: 800%

The bitumen/polymer binder was emulsified by making use either of the procedure of example 6 according to the invention (test 8.I) or of the procedure of the control Example 2 (test 8.II)

In each case there were obtained 1000 parts of a cationic emulsion of the bitumen/polymer binder having the characteristics stated herebelow:

test	8.I	8.II
pH	5.4	3.7
medium diameter of the globules of binder (μ m)	4-7	3-6
rupture index	60	63
quantity of liquid salted out after 1 hour (% of the emulsion)	7	33
quantity of water in the salted out liquid (%)	95	32
cohesiveness of the clot	strong elasticity	zero
time of rupture of the emulsion on plates of coated product (mn)	30-60	> 180

In each one of the tests all of the binder was emulsified during the preparation of the emulsion and the stability in storage of the emulsion produced is good.

For the emulsion of test 8.I according to the invention the rupture index of the emulsion is low and the rupture of the emulsion is accompanied by salting out of reduced amount of liquid containing very little of non-ruptured emulsion. Besides, the clot of sand and bitumen/polymer binder obtained in the course of the test for determining the rupture index has a strong elasticity.

The traction characteristics of this clot obtained by traction test following the standard NF T 46002 working at 0° C. with a traction speed of 500 mm/mn are the following:

Stress at threshold: 3.05×10^5 Pa

Stress at breakage: 0.38×10^5 Pa

Elongation at threshold: 15%

Elongation at breakage: 180%

For the control emulsion of test 8.II, the rupture index has a value approximate to that of the emulsion of test 8.I, but the rupture of the emulsion is accompanied by the salting out of a considerable amount of liquid containing a great proportion of non-ruptured emulsion. In addition, the clot of sand and bitumen/polymer binder obtained in the course of the test for determining the rupture index has no cohesiveness at all. Finally, the time of rupture of the emulsion on plates of coated products is much longer than that measured for the emulsion of test 8.I according to the invention.

EXAMPLE 9

A bituminous binder of the bitumen/polymer type was prepared by mixing 80 parts of a road bitumen of penetration 80/100 brought to 180° C. with 20 parts of a vulcanized mother solution formed from a hydrocarbon solvent, a block copolymer of styrene and butadiene and sulfur as indicated in French Pat. No. 7,910,987 (Publication No. 2,455,623) of May 2, 1979.

The bitumen/polymer binder obtained had the following characteristics:

viscosity at 160° C.: 60 mPa.s

pseudo-viscosity at 50° C. with an orifice of 10 mm (NF T 66 005): 78 seconds
 traction test at -10° C. with a speed of 500 mm/mn
 Stress at threshold: $2,55 \times 10^5$ Pa
 Stress at breakage: $0,14 \times 10^5$ Pa
 Stress at threshold: $2,55 \times 10^5$ Pa
 Stress at breakage: $0,14 \times 10^5$ Pa
 Elongation at threshold: 15%
 Elongation at breakage: >900%

From the bitumen/polymer binder thus obtained, there were prepared two cationic emulsions by making use either of the procedure of example 6 (test 9.I) or of the procedure of the control Example 2 (test 9.II) In each case there were obtained 1000 parts of a cationic emulsion of the bituminous binder of the bitumen/polymer type having the following characteristics:

test	9.I	9.II
pH	5.5	3.8
medium diameter of the globules of binder (μ m)	3-6	3-6
rupture index	57	58
quantity of liquid salted out after 1 hour (% of the emulsion)	6	37
quantity of water in the liquid salted out (%)	97	32
cohesiveness of the clot	good	zero
	cohesiveness	
time of rupture of the emulsion on plates of coated product (mn)	30-60	>180

In each one of the tests all of the binder is emulsified during the preparation of the emulsion and the stability in storage of the emulsion obtained is good.

For the emulsion of test 9.I according to the invention the rupture index of the emulsion is low and the rupture of the emulsion is accompanied by salting out of a reduced amount of liquid containing very little of non-ruptured emulsion. Additionally, the clot of sand and bitumen/polymer binder obtained in the course of the test for determining the rupture index has a good cohesiveness.

The traction characteristics of this clot obtained by traction test according to standard NF T 46002, working at 10° C. with a traction speed of 500 mm/mn are the following:

Stress at threshold: $3,3 \times 10^5$ Pa
 Stress at breakage: $0,27 \times 10^5$ Pa
 Elongation at threshold: 10%
 Elongation at breakage: 230%

For the control emulsion of test 9.II, the rupture index has a value comparable to that of the emulsion of test 9.I, but the rupture of the emulsion is accompanied by salting out of a considerable amount of liquid containing a substantial proportion of non-ruptured emulsion. Besides, the clot of sand and bitumen/polymer binder obtained in the course of the test for determining the rupture index has no cohesiveness at all. Finally, the time of rupture of the emulsion on plates of coated product is much longer than that measured for the emulsion of test 9.I according to the invention.

EXAMPLE 10

From the bitumen/polymer binder obtained as described in Example 2, there was prepared a cationic emulsion according to the invention with 70% of said binder following the mode of operation defined in Example 5, but with two modifications, namely, the use by way of emulsifying agent of a mixture of dimethyl ben-

zyl ammonium tallow chloride sold under the name of NORAMIUM S 75 by the firm CECA and of tallow polypropylene polyamine (LILAMULS EM 30) in respective quantities representing 0.17% and 0.1% of the aqueous phase, and the use of an amount of hydrochloric acid representing only 0.1% of the aqueous phase.

There were obtained 1000 parts of cationic emulsion having the following characteristics:

pH: 5.6
 medium diameter of the globules of binder (μ m): 3-6
 rupture index: 50
 quantity of liquid salted out after 1 hour (% of the emulsion): 10
 quantity of water in the salted out liquid (%): 98
 cohesiveness of the clot: great elasticity
 time of rupture of the emulsion on plates of coated product (mn): 30-60

All of the bitumen/polymer binder was emulsified and the stability in storage of the emulsion produced is good.

The rupture index of the emulsion is low and there is found a behavior of the emulsion similar to that of an ordinary bitumen, namely, small amounts of salted out liquid phase and little of non-ruptured emulsion.

Besides, the clot of sand and bitumen/polymer binder obtained in the course of the test for determining the rupture index has a strong cohesiveness-elasticity characterized by traction test.

EXAMPLE 11

From the bitumen/polymer binder obtained as described in Example 2, there was prepared a cationic emulsion according to the invention with 70% of said binder as follows.

There was first formed a mixture of 45 parts of dimethyl benzyl ammonium tallow chloride (NORAMIUM S 75), 25 parts of tallow polypropylene polyamine (LILAMULS EM 30), 25 parts of an aqueous solution with 40% tetrapotassium salts of hydroxypropane-1, 1-diphosphonic acid and 67 parts of an aqueous solution with 30% of a thickening agent consisting of arabic gum.

An aqueous phase was prepared by incorporating 0.4 part of the mixture thus produced into 100 parts of city water and then adding to the dispersion 0.1 part of hydrochloric acid at 20° Be.

The emulsion was then produced in an emulsifier in which were simultaneously and separately introduced 300 parts of the aqueous phase prepared as indicated above and brought to 45° C. and 700 parts of the bitumen/polymer binder brought to 145° C.

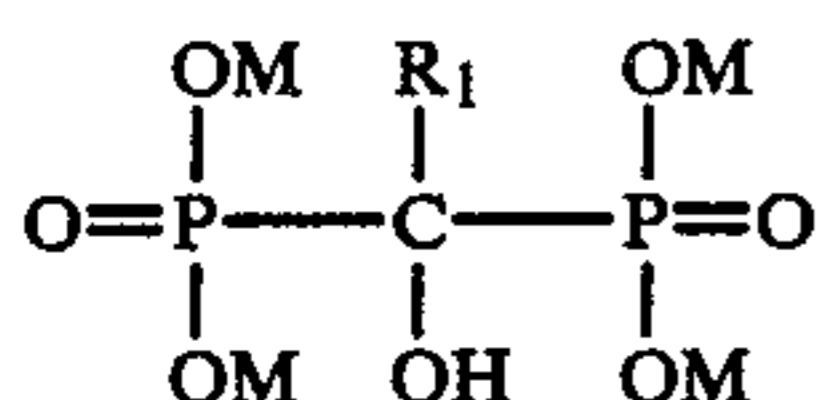
The cationic emulsion obtained had characteristics similar to those of the cationic emulsion obtained in example 10 although its viscosity was higher, namely, above 100 centistokes.

All of the bitumen/polymer binder was emulsified and the stability in storage of the emulsion produced was good.

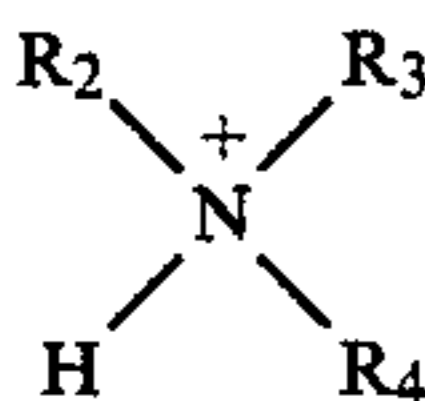
What is claimed is:

1. Cationic emulsions of bituminous binders of the bitumen/polymer type, which are formed from a dispersion of an organic phase comprised of a bitumen/polymer binder in an aqueous phase containing a cationic nitrogen containing emulsifying agent in an amount representing from about 0.03% to 3% by weight of the emulsion and an acid, said acid being present in a quantity such that the cationic emulsifying

agent is rendered water-soluble and the pH of said aqueous phase has a value of from about 3 to 9; wherein said aqueous phase contains an adjuvant consisting of at least one water soluble sequestering salt of phosphonic acid in which the sequestration power in respect to the metallic ions is equivalent, in a pH range of from about 3 to about 9, to a fixation of at least 5 mg. calcium ions per gram of salt, said salt having the formula



wherein R_1 is a hydrocarbyl radical of from C_1 to C_{10} and M , which can be identical or different, designates a hydrogen atom or a group M_1 ; at least one M group is an M_1 group, said M_1 being selected from the group consisting of an alkaline metal atom, $-\text{N}^+\text{H}_2-\text{NH}_2$, and a radical of the formula



wherein the R_2 to R_4 symbols can be identical or different, and each represents one hydrogen atom or a nonvalent hydrocarbyl radical from C_1 to C_8 which can contain one or more functional groups, and wherein two of the radicals R_2 to R_4 can combine to form a heterocyclic group with the nitrogen atom to which they are bonded, the concentration of the adjuvant in the aqueous phase of the emulsion being from about 0.005% to about 0.5% by weight of the emulsion; wherein said emulsion contains by weight from about 30% to about 85% of bitumen/polymer binder; wherein the cationic emulsifying agent present in the emulsion is formed by association of one or more emulsifying agents A selected from the group consisting of quaternary ammonium salts with one or more emulsifying agents B selected among the cationic emulsifying agents having in their molecule, at least three functional groups selected among the group amines and amides in a manner such that at least one of said functional groups is an amine group, and wherein the weight ratio of the total quantity of the compound or compounds A to the total quantity of A and B compounds is from about 5% to about 95%.

2. Emulsions according to claim 1 wherein said R_2 - R_4 are hydrocarbyl radicals of from C_1 to C_8 and said alkaline metal atom is sodium or potassium.

3. Emulsions according to claim 1 wherein the R_1 radical of the formula defining the adjuvant is an alkyl, aryl, or cycloalkyl radical.

4. Emulsions according to claim 1 wherein the hydrocarbyl radicals represented by the symbols R_2 to R_4 are alkyl or hydroxyalkyl radicals from C_1 to C_8 , some of said radicals R_2 to R_4 being capable of forming an aromatic, or alicyclic, heterocyclic group with the nitrogen atom to which they are bonded.

5. Emulsions according to claim 4 wherein two of said R_2 - R_4 radicals form a pyridine or piperidine group with the nitrogen atom to which they are bonded.

6. Emulsions according to claim 1 wherein the concentration of said phosphonic adjuvant is from about 0.01% to about 0.3% by weight of the emulsion.

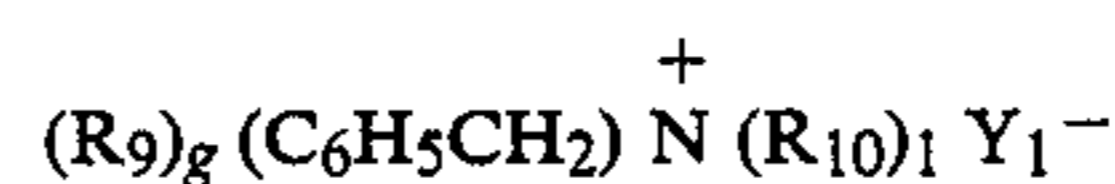
7. Emulsions according to claim 1 wherein the emulsifying agents A are quaternary ammonium compounds of the formula $(\text{R}_7)_j\text{N}^+(\text{R}_8)_h\text{Y}^-$ while the emulsifying agents B are selected among the polyamines and amidopolyamines of the formula



wherein R represents a hydrocarbyl radical, having up to 22 carbon atoms, and m and n are identical or different, being integers from 1 to 8 r being equal to 0 or 1, s representing an integer from 1 to 6, the R_7 's can be identical or different, and designate C_6 to C_{22} hydrocarbyl radicals which can be substituted with one or more functional groups, the R_8 's can be identical or different, representing alkyl or hydrocarbyl radicals from C_1 to C_6 , Y^- designating an anion of a mineral acid or of an organic acid, h being a number equal to $(4-j)$ and j representing a number having the value of 1, 2 or 3.

8. Emulsions according to claim 7 wherein R is alkyl or alkenyl of from C_8 to C_{22} ; and m and n are integers of from 1 to 6; s represents an integer of from 1 to 3; R_7 represents alkyl or alkenyl of from C_8 to C_{22} ; and Y^- represents a chloride, acetate or format anion.

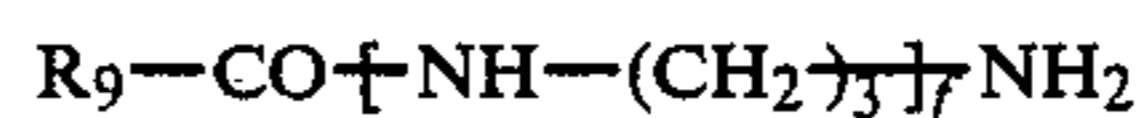
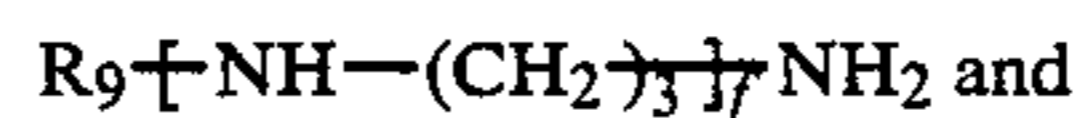
9. Emulsions according to claim 1 wherein the emulsifying agents A are selected among the compounds of formulae:



wherein R_9 designates an alkyl or alkenyl radical from C_{12} to C_{22} , R_{10} represents an alkyl or hydroxyalkyl from C_1 to C_3 , Y_1^- is a chloride or acetate anion, 1 is a number equal to $(3-g)$, g is a number having the values 1, 2 or 3, h is a number equal to $(4-j)$ and j designates a number having the value of 1, 2 or 3.

10. Emulsions according to claim 9 wherein R_9 represents stearyl or oleyl.

11. Emulsions according to claim 1 wherein the emulsifying agents are selected from among the compounds of the formula:



wherein f designates a number assuming the values 2, 3, 4 or 5 and R_9 represents an alkyl or alkenyl radical from C_{12} to C_{22} .

12. Emulsions according to claim 10 wherein R_9 represents stearyl or oleyl.

13. Emulsions according to claim 1 wherein the total quantity of cationic emulsifying agent is from 0.06% to 2% by weight of the emulsion.

14. Emulsions according to claim 1 wherein the acid used to control the pH of the emulsion is a mineral acid selected from the group consisting of hydrochloric, nitric or phosphoric, or a saturated or unsaturated mono or polycarboxylic acid.

15. Emulsions according to claim 13 wherein said acid is selected from the group consisting of acetic, formic, oxalic and citric.

16. Emulsions according to claim 1 wherein the bitumen/polymer binder is selected from the group consisting of bitumens modified by at least one block copolymer of styrene with a diene selected from the group consisting of butadiene, isoprene and a carboxylated diene, and bitumens modified by being put into contact with a mother solution containing at least one styrene/-conjugated diene block copolymer, sulfur or a polysulfide, and a fraction selected from the group consisting of a petroleum fraction, a coal oil, and a mixture of a petroleum fraction, a coal oil, and a mixture of a petroleum reaction and a coal oil.

17. Emulsions according to claim 1 wherein said emulsions also contain, in their aqueous phase, a thickening agent consisting of one or more water-soluble organic compounds, capable within a pH range of from 3 to 9, of increasing the viscosity of the emulsion without modifying the viscosity of the aqueous phase alone, said thickening agent being used in an amount representing from 5% to 30% of the total weight of the combination formed by the cationic emulsifying agent, the phosphoric adjuvant, and the thickening agent.

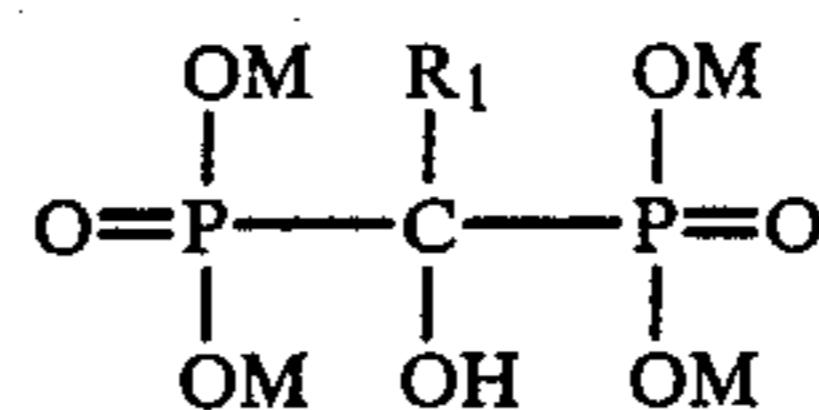
18. Emulsions according to claim 17 wherein the thickening agent is a water-soluble natural gum.

19. Emulsions according to claim 17 wherein the thickening agent is a water-soluble polyurethane of low molecular weight which results from the reaction of a polyisocyanate with a polyester polyol or a polyether polyol.

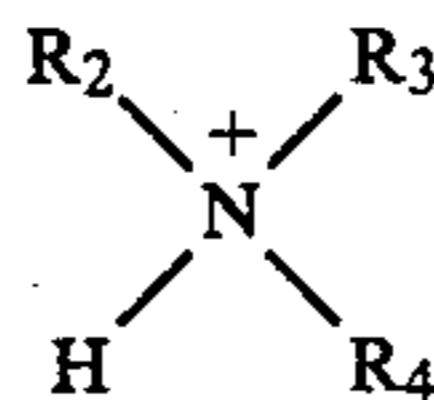
20. Emulsions according to claim 19 wherein the polyurethane has a molecular weight of less than 20000.

21. Cationic emulsions of bituminous binders of the bitumen/polymer type, which are formed from a dispersion of an organic phase comprised of a bitumen/polymer binder in an aqueous phase containing a cationic nitrogen containing emulsifying agent in an amount representing from about 0.03% to 3% by weight of the emulsion and an acid, said acid being present in a quantity such that the cationic emulsifying agent is rendered water-soluble and the pH of said aqueous phase has a value of from about 3 to 9; wherein said aqueous phase contains an adjuvant consisting of at least

one water soluble sequestering salt of phosphoric acid, with the formula



wherein R₁ is a hydrocarbyl radical of from C₁ to C₁₀ and M, which can be identical or different, designates a hydrogen atom or a group M₁; at least one M group is an M₁ group, said M₁ being selected from the group consisting of an alkaline metal atom, —NH₂, —NH₃, or a radical of the formula



wherein the R₂ to R₄ symbols can be identical or different, as each represents one hydrogen atom or a monovalent hydrocarbyl radical from C₁ to C₈ which can contain one or more functional groups, and wherein some of the radicals R₂ to R₄ can combine to form a heterocyclic group with the nitrogen atom to which they are bonded, the concentration of the adjuvant in the aqueous phase of the emulsion being from about 0.005% to about 0.5% by weight 30% to about 85% of bitumen/polymer binder; wherein the cationic emulsifying agent present in the emulsion is formed by association of one or more emulsifying agents A selected from the group consisting of quaternary ammonium salts with one or more emulsifying agents B selected among the cationic emulsifying agents having in their molecule, at least three functional groups selected among the group amines and amides in a manner such that at least one of said functional groups is an amine group, and wherein the weight ratio of the total quantity of the compound or compounds A to the total quantity of A and B compounds is from about 5% to about 95%.

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UNITED STATES PATENT AND TRADEMARK OFFICE
CERTIFICATE OF CORRECTION

PATENT NO. : 4,879,326
DATED : November 7, 1989
INVENTOR(S) : Demangeon, et al

Page 1 of 3

It is certified that error appears in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

- Column 2, line 64: the number of the patent has been omitted and should read --7729953--.
line 66: the word "bi-tumens" should be --bitumens--
- Column 5, line 43: The percentage "0.05 %" should read --0.5 %--.
- Column 6, line 47: the word "amidodiamines" should be --amidoamines--.
- Column 7, line 14: correct the spelling of "formate"
line 33: insert "R₉ being" between "formulae" and "an alkyl or...".
line 42: correct the spelling of "following".
- Column 9, line 13: change "for bring" into --for bringing--.
line 23: the term "alifiying" should read --salifying--.
- Column 10, line 55: correct the spelling of "microdiorite".
- Column 11, line 57: the word "saited" should read --salted--.
line 61: the word "amulsion" should be --emulsion--.
- Column 13, line 6: "in" should be --In--.
- Column 14, lines 19-20: "105 Pa" should be --10⁵ Pa--.
line 59: correct the spelling of "elasticity".
- Column 15, line 11: "o" should be --of--.
line 22: "pa" should read --Pa--
line 34: "cut" should be --out--.
- Column 16, line. 2: "pa" should be --Pa--.

UNITED STATES PATENT AND TRADEMARK OFFICE
CERTIFICATE OF CORRECTION

PATENT NO. : 4,879,326

Page 2 of 3

DATED : November 7, 1989

INVENTOR(S) : Demangeon, et al

It is certified that error appears in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

Column 17, line 4: "pa" should be --Pa--.
line 6: to be cancelled.
line 7: to be cancelled.

Column 18, line 16: correct the spelling of "elasticity".

Column 19, line 19: the formula " $-N^+H_2-NH_2$ " should read
-- $-NH_3-NH_2$ --.
line 39: correct the spelling of "present"
line 65: "t" should be --to--

Column 20, line 26: correct the spelling of "formate"
line 38: "y-1" should read --Y₁--.
line 45: insert --B-- between "agents" and "are".

Column 21, line 12: the word "reaction" should read --fraction--.
line 20: "305" should be --30 %--.
line 33: "tupe" should be --type--
line 39: correct the spelling of "emulsifying".

Column 22, line 1: change "phophoric" into --phosphonic--.
line 13: the formula " $-NH_2-NH_2$ " should read
-- $-NH_3-NH_2$ --.
line 22: "as each" should be --and each--.

UNITED STATES PATENT AND TRADEMARK OFFICE
CERTIFICATE OF CORRECTION

PATENT NO. : 4,879,326
DATED : November 7, 1989
INVENTOR(S) : Demangeon, et al

Page 3 of 3

It is certified that error appears in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

Column 22, line 29: between "by weight" and "30 %" should be inserted the following omitted sentence --of the emulsion, wherein said emulsion contains by weight from about--.

Signed and Sealed this
First Day of October, 1991

Attest:

Attesting Officer

HARRY F. MANBECK, JR.

Commissioner of Patents and Trademarks